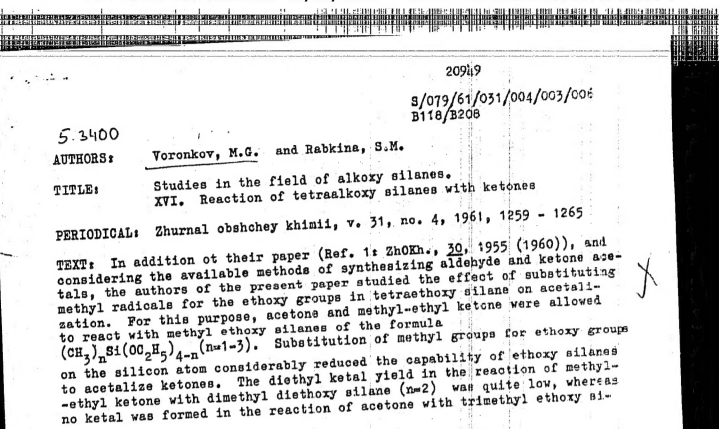
Card 1/6



"APPROVED FOR RELEASE: 03/20/2001 CIA-RDP86-00513R001860920006-9 20919 8/079/61/031/004/003/006 B118/B208 Studies in the field of alkoxy . lane. As acetals in general, and ketals in particular, split off alcohol at elevated temperatures, the synthesized acetals were distilled in the presence of Na or its alcoholate, the acetals otherwise being contaminated by alcohol and the corresponding α,β-unsaturated ether. The ketale of cyclohexanone and its derivatives readily split off alcohol during distillation, and are converted to 1-alkoxy cyclohexenes. This ability of 1,1-dialkoxy cyclohexanes easily yields 1-alkoxy cyclohexenes which are formed immediately from cyclohexanone and tetraalkoxy silaness  $- OR + \frac{1}{m} ((RO)_{4-2n} Sio_n)_m + nROH.$  $=0 + Si(OR)_4 \xrightarrow{H+} n\langle$ Synthesis was performed by simple distillation of a mixture of cyclonexanone and the corresponding tetraslkoxy silane in the presence of H3PO4. This mixture had previously been boiled for 8-14 hr. Rearly all the alcohol is first distilled off, and immediately afterwards the reaction product containing some alcohol. It was purified by distillation on Na, most suitably at low pressure. Five 1-alkoxy cyclohexenes were synthesized in

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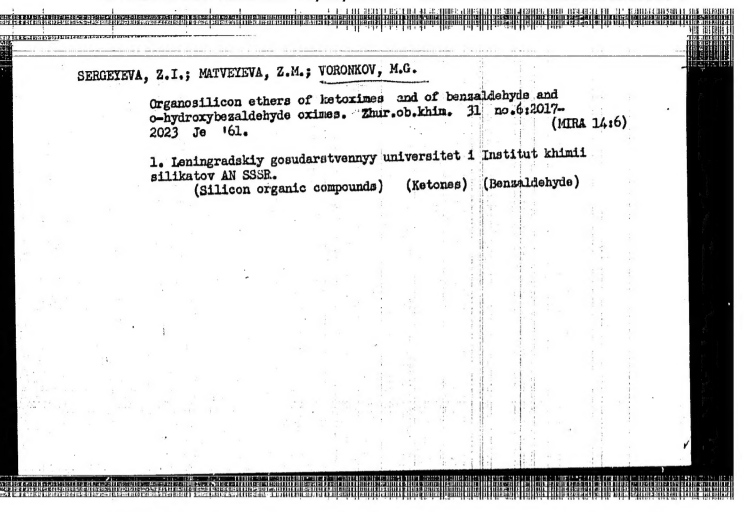
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Studies in the field of alkoxy ..

this way with yields between 55 and 70% (Table). The presence of a certain quantity of alcohol in the reaction product is important in acetalization. It was shown that the reaction of tetraalkoxy silanes with aliphatic and alicyclic ketones takes place in the presence of anhydrous orthophosphoric acid as a catalyst and in the absence of alcohole, forming the corresponding ketals. Eight ketals were obtained and identified in this way, as well as the acetal of butyraldehyde. The opinion expressed in publications that aldehydes are acetalized more easily than ketones may only be explained by the fact that ketals hydrolyze more rapidly than aldehyde acetals. Yu.N. Platonov is thanked for making the analyses. There are 1 figure, 1 table, and 26 references: 14 Soviet-bloc and 12 non-Soviet-bloc. The 2 references to English-language publications read as follows: H.E. Carswell, H. Adkins, J.Am.Chem.Soc., 50, 235, (1928); G.L. Pfeifer, H. Adkins, J.Am.Chem.Soc., 53, 1043 (1931).

ASSOCIATION Institut khimil silikatov Akademii nauk SSSR - (Institute of Silicate Chemistry of the Academy of Sciences USSR)

Card. 3/6



15.6400 1583

22135 S/080/61/034/007/010/016 D223/D305

AUTHORS:

Voronkov, M.G., Sharonov, G.P., and Dolbin, Y.V.

TITLE:

Effect of the nature of sulpho-organic compounds in oil on the frictional wear of metallic surfaces

PERIODICAL: Zhurnal prikladnoy khimii, v. 34, no. 7, 1961,

1562 - 1569

TEXT: The initial wear of new machine parts during the "running in" time can be accelerated by use of sulphurated oils; actual trials have given a time period of 1.5 - 2.0 hours. (Ref. 1: G.P. Sharonov, V.S. Nikandrov, Tankist, 9, 54, 1957). In this connection investigation into the effect of sulphur compounds in oil on running-in and initial machine wear is important both from the theoretical and practical aspects. The article gives the results of investigations on sulphurated oil, its nature, and surveys new sulphurous additions to the oil. The sulphurated oils "industrial 50" and "spindle AV" were produced as follows: To the neated oil

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22435

S/080/61/034/007/010/016 D223/D305

Effect of the nature of ...

in a bath at 130-135°C, slowly and with continuous stirring 4.5 wt. % of sulphur flour was added - the additions taking 2 hours - and gradually raising the oil temperature to 150°C. The oil was kept at this temperature for 24 hours, continuously stirring, the total heating time being 28 hours. The sulphurated oil "industrial 50" was obtained containing 4.23 % of sulphur (Ref. 3: Ye.Ya. Anten, N.V. Mitrofanova, T.N. Abramova, G.P. Sharonov, V.S. Nikandrov, Avt. svid. 20319, 1959). The testing on copper sheets showed the disappearance of corrosion for "spindle AV" after 9 hours of sulphuration and for "industrial 50" after 6 hours. The removal of corrosion by the action of sulphur and with increase in the sulphuration time of oil is probably due to the transition of free sulphur into compounded, as well as the elimination from the oil of H2S and decomposition of mercaptans, since H2S is a product of the reaction of sulphur and hydrocarbons. The friction experiments are then described using machine MI and lubricant MT-16, obtained from sulphurated and emba natural crude oil. These oils had similar viscosities and contained 1.11 and 0.41 % of natural sulphur respec-

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S/080/61/034/007/010/016 D223/D305

Effect of the nature of ...

tively. In addition to emba crude oil, sulphurated oil was added containing 1.11 % of sulphur, of which 0.41 % was natural sulphur. The frictional drum was made of steel 40% having a microhardness 300-310 while the brake was made from steel 45 which after heat treatment showed a microhardness of 660-680. Fig. 2 shows the curves of temperature change of the surface layer of the brake and frictional movement against the working time of steel samples. Since experimental time was 8 hours per day the curves show discontinuity. It follows that pretreatment of samples using emba oil MT-16 is complete in 68 hours (curve 3, Fig. 2). By this time the frictional movement and temperature of the surface layer measured by a thermocouple have reached their minimum values. The pretreatment of samples in emba oil to which sulphurated oil was added containing 1.11 % of sulphur, was complete in 5 hours (curve 1, Fig. 2) and the minimum values of frictional movement and surface layer temperature did not change after an additional 65 hours. Samples lubricated with MT-16 from crude oil with 1.11 % of natural sulphur were not complete in 70 hours (curve 2, Fig. 2) and

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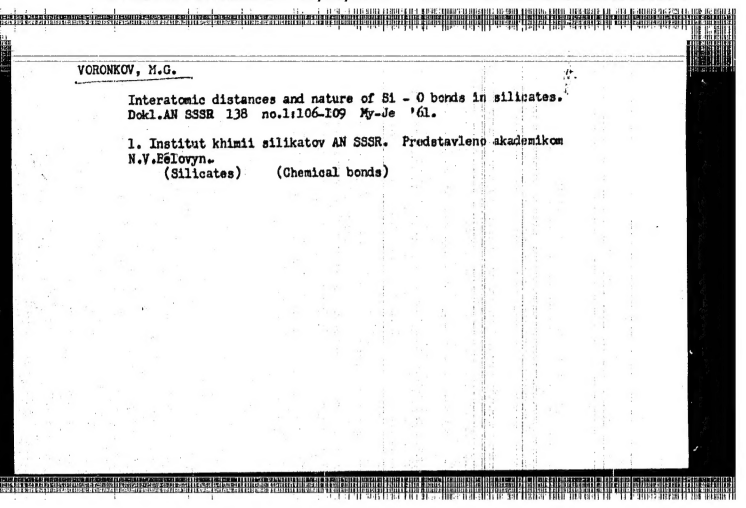
movement and temperature curves show small gradual decrease with time. The contact area of brake to drum was only 35-40 % compared to the sulphurated oil of nearly 100 %. Tests with oils MK with 0.6 % of molybdenum disulphide, AK-10 with additions of different quantities of IP-22 and industrial 12 with 1.5 % of tsiatim-5 did not give positive results, since they behaved in the same manner as natural sulphur. The sulphurated oil protects the working parts of machine by the formation of thin layer of sulphides FeS, FeS2, Fe3S4. With the formation of iron sulphides, the cubical lattice of alpha-Fe changes into hexagonal lattice FeS which, by analogy with graphite, possesses lubricating properties. To investigate the problem of effect of interaction products of S and hydrocarbons and also the additions of different organic compounds of sulphur, tests were done with a series of organic compounds (mercaptan sulphide, de- and polysulphides, thiophen etc.). The results show that disulphides, in particular dibenzencially hide have the same effect as sulphurated oil, while other groups of sulphur organic compounds have not. This necessitated an examination of organic

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Effect of the nature of ... S/080/61/034/007/010/016 D223/D305

polysulphides R<sub>2</sub>Sn, preferably dibenzenepolysulphides o<sub>6</sub>H<sub>5</sub>CH<sub>2</sub>SnCH<sub>2</sub> C<sub>6</sub>H<sub>5</sub> with n >2. These were prepared by adding corresponding quantities of sulphur to dibenzenedisulphide at 150°C. These compounds with 0.9 - 1.0 % of sulphur were found to be quite effective. There are 5 figures and 3 references: 2 Soviet-bloc and 1 non-Soviet-bloc. The reference to the English-language publication reads as follows: Diesel power, 35, 11, 32, 1957.

SUBMITTED: February 1, 1961



# VORDNKOV, MIKHAIL GRIGOR VEVICH

PHASE I BOOK EXPLOITATION

SOV/6267

Davydova, Viktoriya Pavlovna, and Mikhail Grigor'yevich Voronkov

Polifosfazeny; polimernyye i monomernyye fosfonitril'nyye soyedineniya (Polyphosphazenes; Polymeric and Monomeric Phosphonitrile Compounds) Moskva, Izd-vo AN SSSR, 1962. 86 p. 2600 copies printed.

Sponsoring Agency: Akademiya nauk SSSR. Institut khimit silikatov.

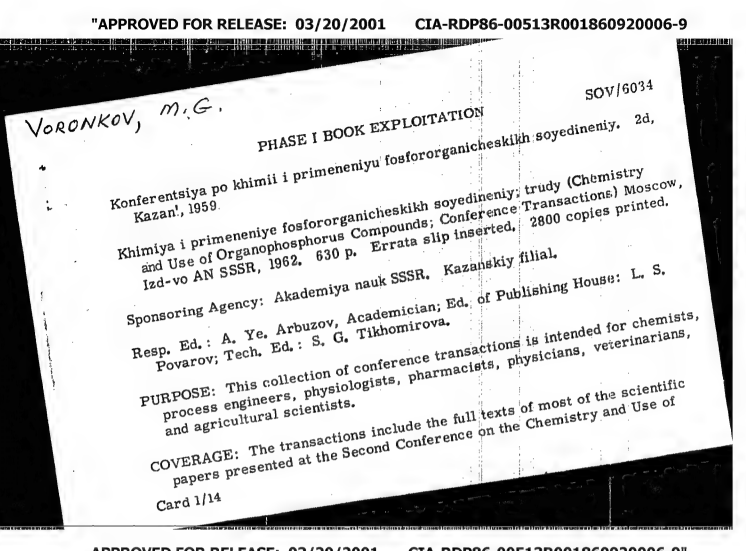
Resp. Ed.: M. G. Voronkov, Doctor of Chemical Sciences; Ed. of Publishing House: V. D. Piastro; Tech. Ed.: G. P. Aref'yeva.

PURPOSE: This book is intended for scientists, engineers, aspirants, teachers, and students in advanced courses at schools of higher education working in the field of monomeric and high-molecular compounds.

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APPROVED FOR RELEASE: 03/20/2001 CIA-RDP86-00513R001860920006-9"

Polyphosphazenes (Cont.	<b>)</b>					
					V/6267	
COVERAGE: This book the existing literature	is reportedly the f	irst effort	to review	and sys	tematize	
the existing literatur and patents published	up to 1961. It is	expected t	hat maten	forter many		
month buoghnour this	polymers will be l	hetter than	matanial	France Al	la a la la M	
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Chemistry and the Use of Organophosphorus (Cont.)

SOV/6034

Organophosphorus Compounds held at Kazan' from 2 Nov through 1 Dec 1959. The material is divided into three sections: Chemistry, containing 67 articles; Physiological Activity of Organophosphorus Compounds, containing 26 articles; and Plant Protection, containing 12 articles. The reports reflect the strong interest of Soviet scientists in the chemistry and application of organophosphorus compounds. References accompany individual reports. Short summaries of some of the listed reports have been made and are given below.

TABLE OF CONTENTS:[Abridged]:

Introduction (Academician A. Ye. Arbuzov)

3

## TRANSACTIONS OF THE CHEMISTRY SECTION

Gefter, Ye. L. [NII plastmass (Scientific Research Institute of Plastics, Moscow]. Some Prospects for the Industrial Use of Organophosphorus Compounds

46

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Chemistry and the Use of Organophosphorus (Cont.)

SOV/6034

The preparation and uses of organophosphorus compounds are discussed, and prospects for obtaining new high-molecular organophosphorus compounds are outlined. Gefter recommends 1) the preparation of new organophosphorus polymers based on oxides of unsaturated tertiary phosphines, 2) the utilization of thermal stability and good mechanical properties of known organophosphorus compounds as a basis for creating new heterochain phosphorus-containing compounds, and 3) application of the latest methods currently used in the chemistry of high-molecular compounds to organophosphorus

Orlov, N. F., and M. G. Voronkov [Institut khimii silikatov (Institute of Silicate Chemistry, Leningrad)]. New Methods of Synthesis of Triorganosilyl Esters of Orthophosphoric and Alkylphosphonic Acids New methods have been developed for the synthesis of tris-(triorganosilyl)-phosphates (R, SiO), PO, bis-(triorganosilyl)methylphosphonates CH3 (R3SiO)2PO, and organophosphorussilicon polymers based on H<sub>3</sub>PO<sub>4</sub> and CH<sub>3</sub>PO(OH)<sub>2</sub>. One method is based on the interaction of

212

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ORLOV, N.F., VORONKOV, M.G.

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New ways to synthesinze triorganceilyl esters of orthophosphoric and alkylphosphinic acids.

Collection of complete papers presented at the 19 5 Mazan Jun'erence us Chemistry of expanophosphorus Commounts.

33988 s/062/62/000/002/013/013 B117/B138

5.2420 5.2410 AUTHORS:

Shchukovskaya, L. L., Voronkov, M. G., and Pavlova, O. V.

TITLE:

New method of N-dimethyl-B-difluoro borazene synthesis

PERIODICAL:

Akademiya nauk SSSR. Izvestiya. Otdeleniye khimicheskikh

nauk, no. 2, 1962, 366

TEXT: The new method consists in the separation of hydrogen fluoride from N-dimethyl-B-trifluoro borazane heated with aluminum dust in vaseline oil medium. C<sub>2</sub>H<sub>7</sub>NBF<sub>3</sub> (boiling point 149 - 150°C (5 mm Hg), dry aluminum dust, and vaseline oil (boiling point 210 - 230°C (2 mm Hg)) were slowly heated and vaseline oil (boiling point 210 - 230°C (2 mm Hg)) were slowly heated in a distilling flask on Wcod's alloy. At 278° hydrogen started separating in a distilling flask on Wcod's alloy. At 278° hydrogen started separating energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way be nearly and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way energetically and the collecting vessel connected with the gasometer by way ener

New method of N-dimethyl-B-difluoro...

33988 S/062/62/000/002/013/013 B117/B138

manner. Hydrogen fluoride could not be separated from N-dimethyl-B-tri-fluoro borazane by heating with KF. There are 2 non-Soviet references. The two references to English-language publications read as follows: J. F. Brown, J. Amer. Chem. Soc. 74, 1219 (1952); A. B. Burg, J. Banus, J. Amer. Chem. Soc. 76, 3903 (1954).

ASSOCIATION: Institut khimii silikatov Akademii nauk SSSR (Institute of Silicate Chemistry of the Academy of Sciences USSR)

SUBMITTED: August 10, 1961

Card 2/2

33923

S/079/62/032/002/008/011 D204/D303

5.3700

. . .

Orlov, N.F. and Voronkov, M.G.

TITLE:

AUTHORS:

Silico-organic esters of methyl phosphinic acid (MPA)

PERIODICAL:

Zhurnal obshchey khimii, v.32, no. 2, 1962, 608-612

TEXT: This is a description of the methods of synthesis of the hitherto unknown bis (triorganosily1) methyl phosphinates, based on interaction of  $\mathrm{CH_3P(0)(OH)_2}$  with: (1) Triorganosilanols:  $\mathrm{CH_3P(0)/OSEEt_3}$  was prepared in 68% yield from  $\mathrm{Et_3SiOH}$  (2 moles) and MPA (1 mole), by heating for 2 hrs, removing continuously the water formed. (2) Triorganosilanes:  $\mathrm{CH_3P(0)}$   $\mathrm{COSIEt_2Me}$   $\mathrm{I}_2$ ,  $\mathrm{CH_3P(0)}$   $\mathrm{I}_2$   $\mathrm{I}_3$   $\mathrm{I}_2$ ,  $\mathrm{CH_3P(0)}$   $\mathrm{I}_3$   $\mathrm{I}_3$ 

Card 1/3

33923

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Silico-organic esters of ...

rSi-0-Si-0-P-0,
Et Et 0
n
was obtained from an equimolecu.

whilst another polymer:  $\begin{pmatrix} \text{CII} & \text{CH}_3 \\ \text{-0-P-0-Si-} \\ \text{0} & \text{CH}_3 \end{pmatrix}$ 

lar mixture of MPA and dimethyl diethoxysilane. All the phosphinates, with the exception of crystalline CH<sub>3</sub>P(0) [ OSi(Ph)<sub>3</sub>]<sub>20</sub> were colorless, high-

boiling liquids, readily hydrolyzed by water. The polymers were pale yellow oils. Experimental details are given and physical constants of

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33923 S/079/62/032/002/008/011 D204/D303

Silico-organic esters of ...

the products are tabulated. There are 1 table and 17 references: 11 Soviet-bloc and 6 non-Soviet-bloc. The 4 most recent references to the English-language publications read as follows: R.O. Sauer, J.Am.Chem.Soc., 66, 1707, (1944); W.H. Keeber and H.W. Post, J.Org.Ch., 21, 509, (1958); Canadian Pat. 496,623 (1953); Ch.A., 47, 4128, (1953).

ASSOCIATION:

Institut khimii silikatov Akademii nauk SSSR (Institute of

Silicate Chemistry of the Academy of Sciences, USSR)

SUBMITTED:

March 2, 1961

Card 3/3

#### "APPROVED FOR RELEASE: 03/20/2001 CIA

CIA-RDP86-00513R001860920006-9

S/079/62/032/002/009/011 D204/D303

AUTHORS:

Orlov, N.F., Bogatkin, R.A., Sergsyeva, Z.I., and Veronkov,

M.G.

TITLE:

Interaction of triorganosilanes with carboxylic acids in

the presence of colloidal nickel

PERIODICAL:

Zhurnal obshchey khimii, v. 32, no. 2, 1962, 650-651

TEXT: A short note on the reactions of triethyl silane with carboxylic acid, using colloidal Ni as a catalyst. Monocarboxylic acids reacted giving the corresponding triorganosilyl esthers, in 50-85% yield. Esters of general formula Et SiOCO(CH<sub>2</sub>) OCOSiEt were synthesized in 60-80% of general formula Et SiOCO(CH<sub>2</sub>) OCOSiEt were synthesized in 60-80%

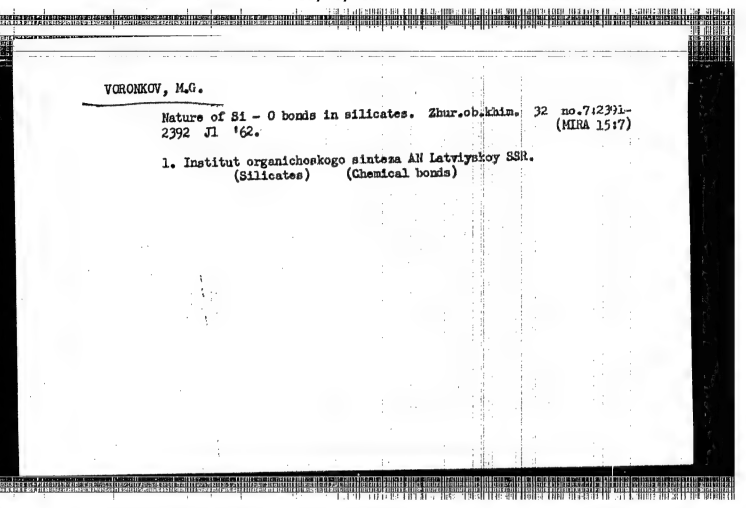
yields from simple dicarboxylic acids. Colloidal Ni promoted hydrogenation as well as dehydrocondensation, as was shown by the reactions of Et\_SiH with halogenated and unsaturated acids. Monochloracetic acid yielded either Et\_SiOCOCH\_Cl or (Et\_SiOCOCH\_3 + Et\_SiCl), depending on the molar ratio of the reagents. Unsaturated acids yield hydrogenated

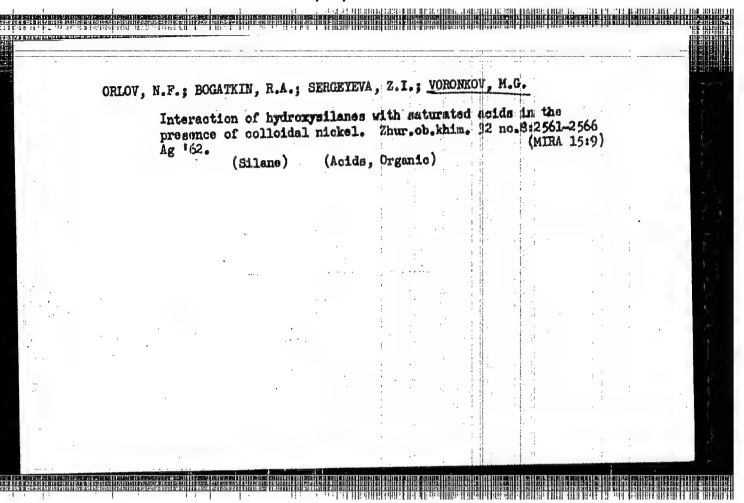
Card 1/2

S/079/62/032/002/009/011
Interaction of triorganosilanes ... D204/D303

products; e.g. crotonic acid gave the n-butyric ester. Action of H2PtCl6 as the catalyst is similar to that of colloidal Ni. No experimental details are given. There are 3 Soviet-bloc references.

SUBMITTED: July 17, 1961





35695

\$/080/62/035/003/015/024 D227/D302

15.7140

AUTHORS:

Lasskaya, Ye. A. and Voronkov, M. G.

TITLE:

Application of organosilicon compounds for hydrophobing

heat insulating articles made of expanded perlite

Zhurnal prikladnov khimii, v. 35, no. 3, 1962, 612-617 PERIODICAL:

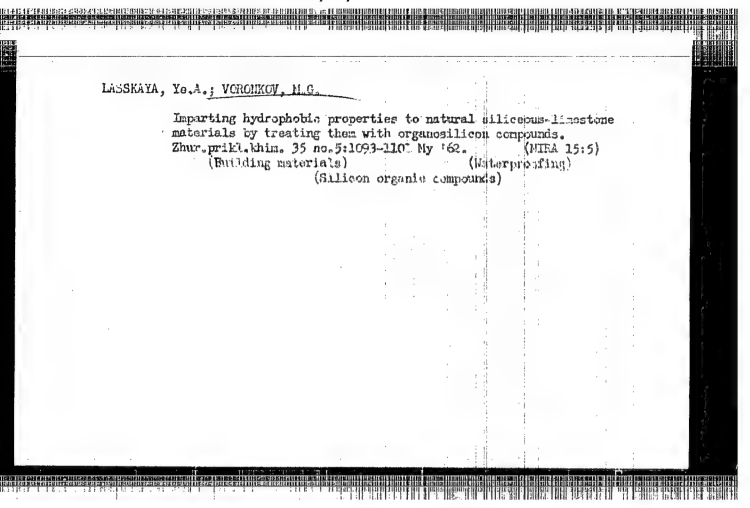
TEXT: To improve water resistance of articles made of expanded perlite, the authors studied the use of organosilicon compounds as possible hydrophobic agents. The compounds tested were: aqueous solution of polyethylhydrosiloxane  $\Gamma K \mathcal{H} - 94$  (GKZh-94) and aqueous solutions of sodium methyl siliconate  $M \supset \Gamma - 9$  (MSG-9) and ethyl siliconate GKZh-10. The solutions were deposited onto air-dried perlite samples in one or two layers and their effectiveness was determined by immersing the samples in water and measuring the amount of water absorbed. The results showed that water-proofing of articles substantially reduces the water absorption capacity. For perlites bonded with concrete, glass and silica, GKZh-94 (~10%) proved to be very effective. The high hydrophobic activity of polyethylhydrosi-Card 1/2

S/080/62/035/003/015/024
Application of organosilicon ... D227/D302

loxanes on the alkali-containing materials is explained by the chemical reaction of the surface materials with Si-H bond giving an organosilicon film and also by hydrolysis of this bond resulting in fusion of the polysiloxane chains. For gypsum and lime-bonded perlites, sodium alkyl silicates (3 - 4%) proved more effective. In general, treatment of various perlites with the mentioned silicon compounds tended to increase their water resistance by up to 65 times. A series of experiments has also been conducted to determine the effect of hydrophobing treatment on other properties of expanded perlite such as water desorption and frost stability. It has been shown that after the treatment with up to 5% solutions of the silicon compound, the samples could withstand up to 25 "freeze-thaw" cycles and up to 30 of "wet-dry" cycles. There are 8 tables and 2 Soviet-bloc references.

SUBMITTED: September 26, 1961

Card 2/2



SHCHUKOVSKAYA, L.L.; VORONKOV, M.C.; PAVLOVA, O.V.

New methods of synthesizing Amonohalo-substituted R-dialkyl-borazines and N-trialkylborazanes. Dokl. AN SSSR 143 no.4; 887-889 Ap '62. (MIRA 15;3)

1. Institut khimii silikatov AN SSSR. Predstavleno akademikom A.V.Topchiyevym. (Borazane) (Borazine)

\$/020/62/145/004/017/024 B110/B144 Lukevits, E. Ya., Romadan, Yu. P., Giller, S. A., Academician AS LatSSR, and Voronkov, M. G. AUTHORS: Organosilicon compounds of the furan series. Organosilicon derivatives of furyl carbinols and 5-substituted furfuryl TITLE: alcohols Doklady, v. 145, no. 4, 1962, 806 - 808 Akademiya nauk SSSR. TEXT: Furfuryl oxysilanes were produced: (1) by reaction of trialkyl chlorosilanes with furyl alkyl and furyl aryl carbinols, (2) by reaction of silanes with furfuryl alcohol, 5-methyl furfuryl alcohol, and furyl alkyl carbinols R = CH4; H, R' = CH4; GH4; CaH4; H R"= CHs ; C1H6 ; C1H6O ; (C1H61251OH R" 15 C4H6 ; C2H7 ; C1H6O using 10-5 moles of H2PtCl6 per 1 mole of isopropyl alcohol as catalyst at Card 1/6 2

3/020/62/145/004/017/024 3110/3144

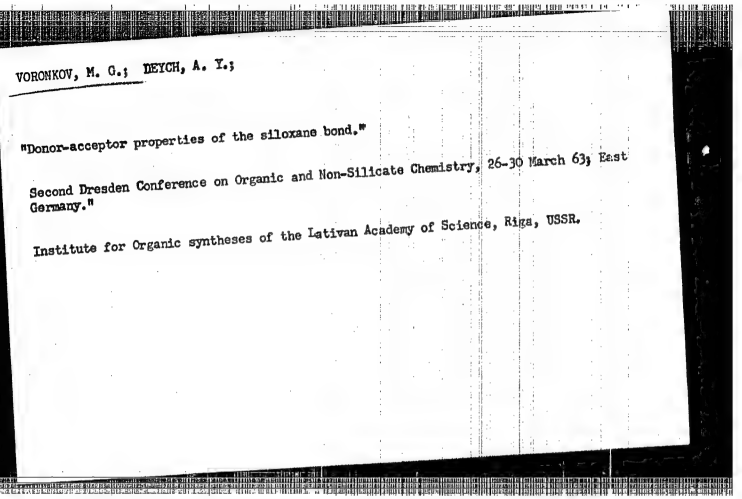
Organosilicon compounds of the ...

80 - 100°C. Their composition and properties are indicated (Table). 5-nitro-furfuryl oxytrimethyl silane was obtained from ethereal solutions of 5-nitro-furfuryl alcohol, pyridine, and trimethyl chlorosilane. Silane reacts with H<sub>2</sub>PtCl<sub>6</sub> in isopropyl alcohol to give furfuryl oxysilane. Dioxane containing 0.05 moles of H<sub>2</sub>O hydrolyzes triethyl silane in the presence of H<sub>2</sub>PtCl<sub>6</sub> to give triethyl silanol. Triethyl silane reacting with triethyl silanol in the presence of H<sub>2</sub>PtCl<sub>6</sub> yields small amounts of hexaethyl disiloxane by anhydrocondensation. There is 1 table.

ASSOCIATION: Institut organicheskogo sinteza Akadenii nauk LatvSSR (Institute of Organic Synthesis of the Academy of Sciences LatSSR)

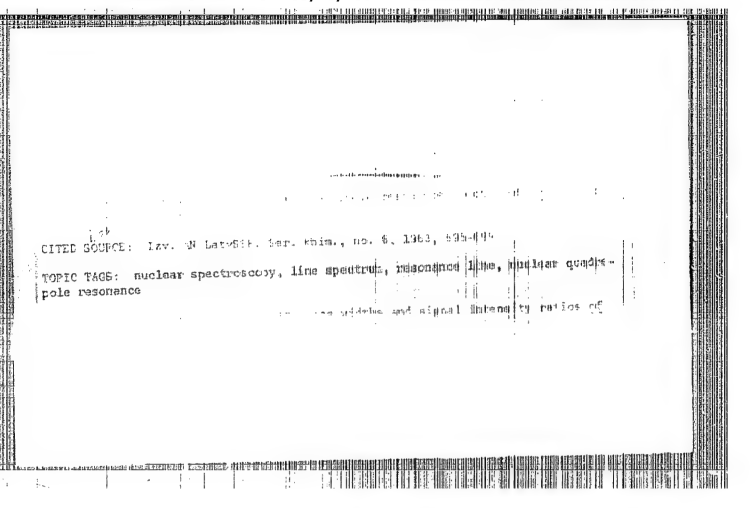
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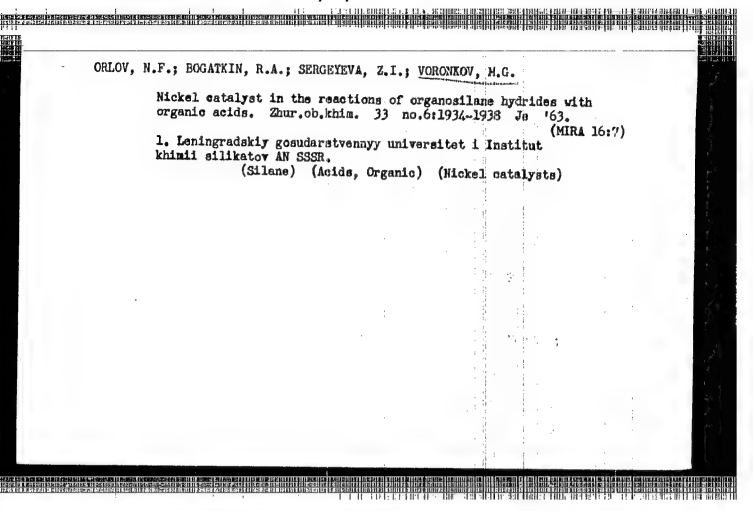
Table. Furfuryl oxysilanes (R'-V-R'). Legend: (1) mode of production, (2) boiling point, °C, (3) pressure, mm Hg.

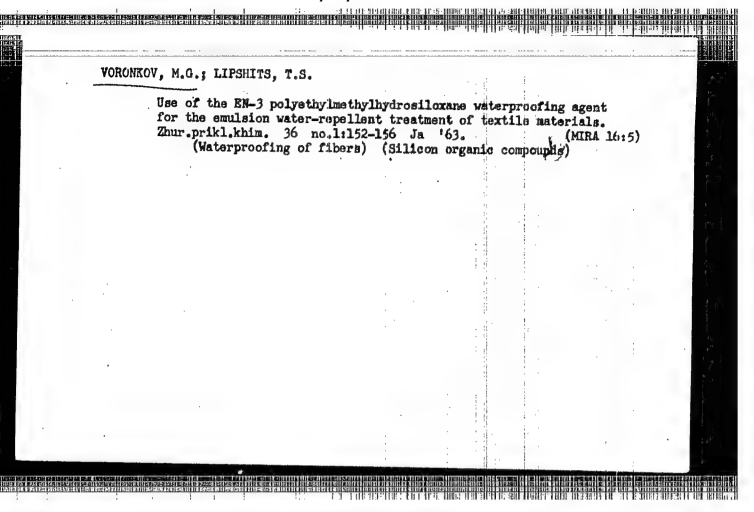


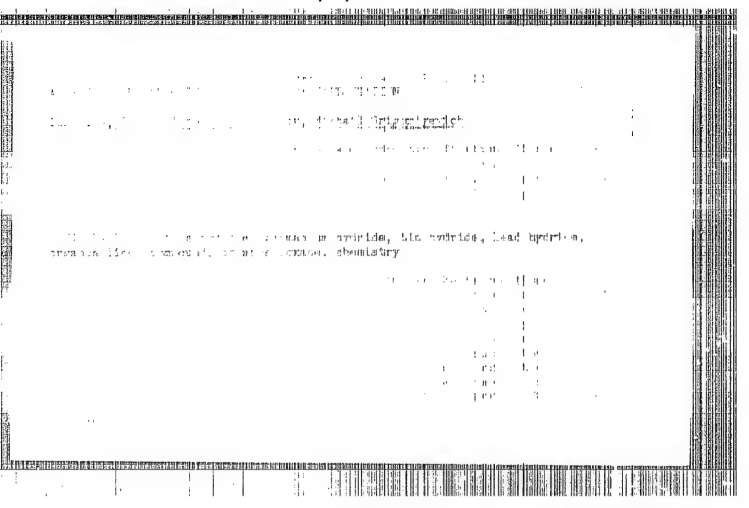
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L 19424-65 ACCESSION NR: AR4048174	
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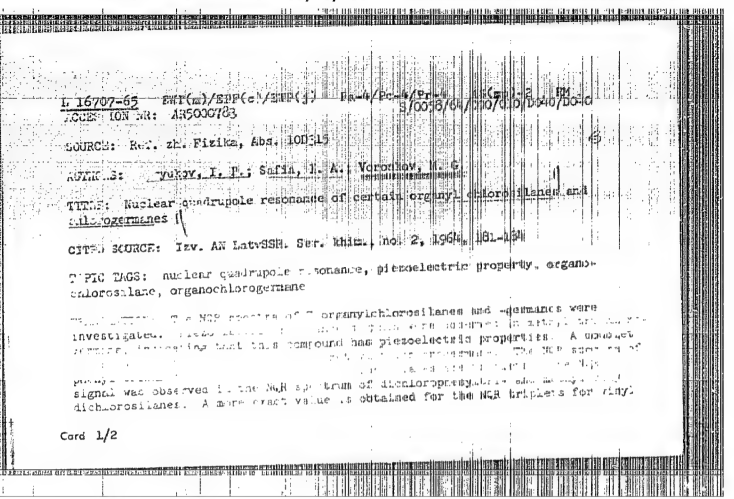
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VORONKOV, M. G.; DEYCH, A. Ya., Rige

"Acceptor complexes of monosubstituted benzenes Compounds."

report submitted for 8th Intl Conf on Coordination Chemistry, Vienna, 7-11 Sep 64,

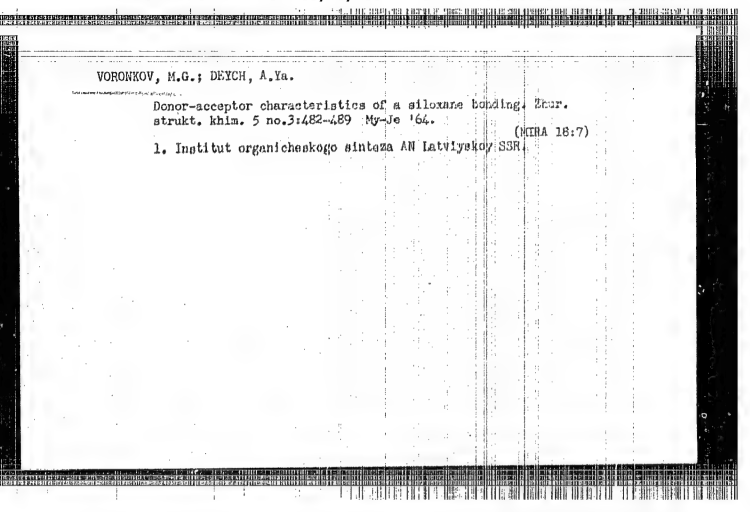


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Park/Prak/Paul SWILL TAR OF TAR THEFTON a Apolice / 6/11/coop/orz / 12 15 / 12/2 . ACCESSION NR: AF4042871 AUTHOR: Toronkov, H. G.; Sharik, Yu. L. Synthesis of organofluorosilanes TITLE: SOURCE: AN SESR. Is westlye. Serira khinichanka, mm. 7, 1951, 1217-1221 TOPTS TAGE: organof morostlace, synthesis, glubrolysis, smmmitum bildhoride. portangings of fluoride story flore stoace cyclinality if happened hand, organized to the si lane, trialhylchic oslime ABSTRACT: A direct and incurrentive method was worked out flow the chuthouts of the state of the second of the second open and the second empydrove powderec githasetin i sammalin baffinomkon RustClook + (2 - 2, ex. M.H.) + RustFa + (2 - 0, b) at HCl + (2 - 0) at a Rus + ) The foliceing alkyl, systemikyl suf arylalkylfluorosilmnos wenu obdadnod in 80-50% yields: trinethyl(whyl on a group)limprodiament die hylfd haythan mitaller on a course and the state of t Card 1/3

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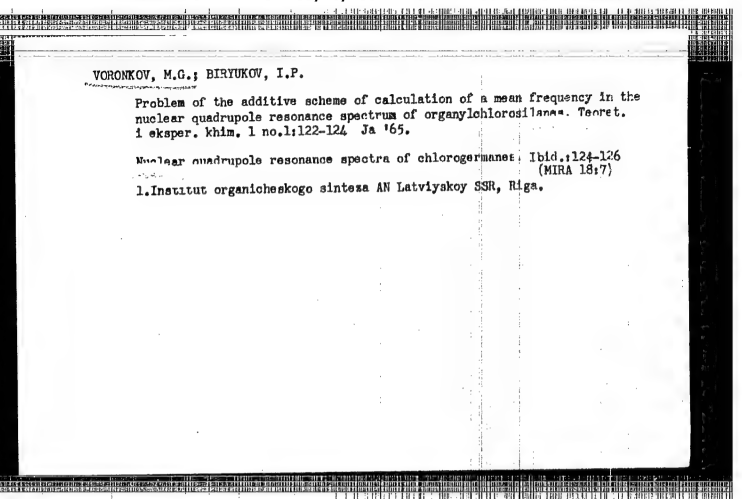
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SOKOLOV, G.P.; GILLER, S.A., akademik; VOROHKOV, M.G.

Reaction of organomagnesium compounds with 2,5-dimethray-2,5-dihydrofurans.
Dokl. AN SSSR 158 no.3:675-678 S \*64. (MIRA 17:10)

1. Institut organicheskogo sinteza AN Latviyskoy SSR. 2. AN Latviyskoy SSR (for Giller).

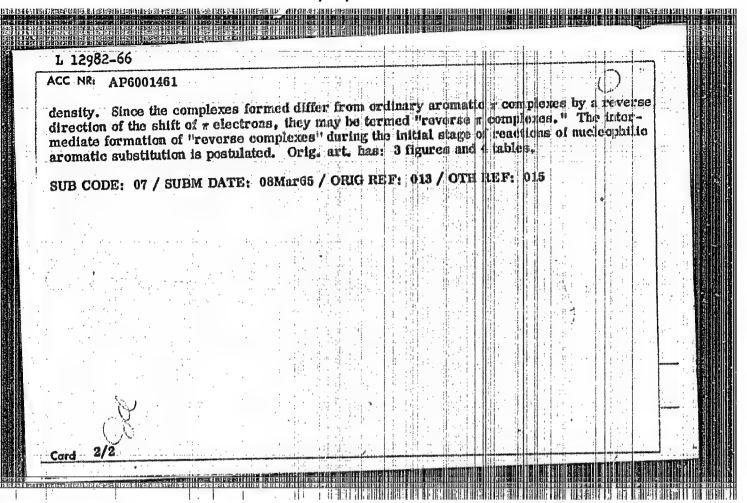


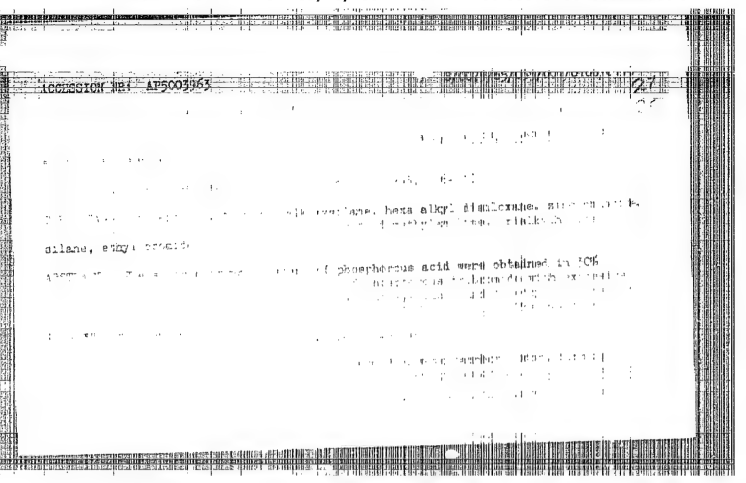
BIRYUKOV, I.P.; VORONKOV, M.G.; SAFIN, I.A.

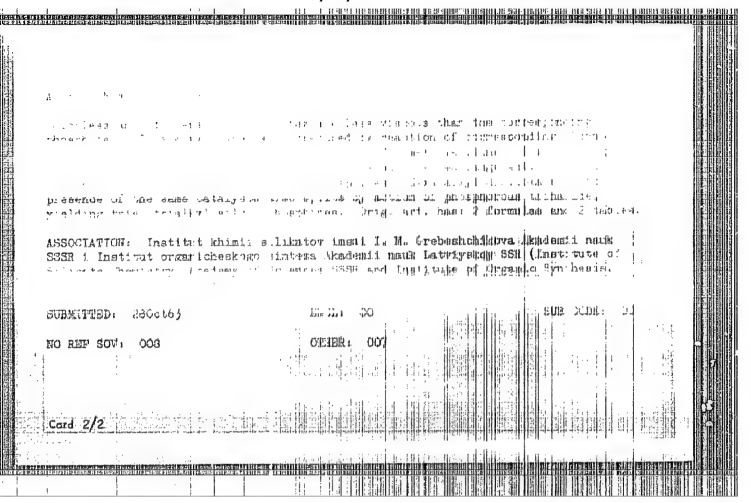
Study of the induction effect of substituents in organochlorosilanes by the nuclear quadrupole resonance method. Touret, 1 eksper. khim. 1 no.32373-380 ky-Je '65. (MIRA 18:9)

1. Institut organicheskogo sinteza AN Latviyskoy SSR, Riga.

ACC NR: AP6001461 SOURCE CODE: UR/0379/65/001/005/0663/0674
AP6001461 SOURCE CODE: UR/0379/05/001/005/0503/067
AUTHOR: Voronkov, M. G.; Deych, A. Ya.
ORG: Institute of Organic Synthesis, AN Latv. ESR, Riga (Institut organic leskogo sinteza); Riga Institute of Civil Aviation Engineers (Hizhskiy Institut inzhendrov granddanskoy aviatsii)
TITLE: Formation of complexes with charge transfer in systems of monosubstituted bearings
SOURCE: Teoreticheskaya i eksperimental'naya khimiya, v. 1, no. 6, 1955, 663-674
10110 1AGS: Intermolecular complex, benzene, electron donor
ABSTRACT: The paper deals with the formation of labile complexes with charge transfer in
that the tendency of C <sub>e</sub> H <sub>5</sub> molecules to form complexes is most property. The authors found orientant, and increase with the found orientant, and increase with the found orientant.
the hetero atom in D: O > Cl > N, and is not proportional to the hetero atom of
component. Complex-formation in $C_0H_0X_0$ systems has a donor-adopton character and is due to the transfer of an electron from D to the aromatic ring with the lower $\pi$ -electron
Card 1/2





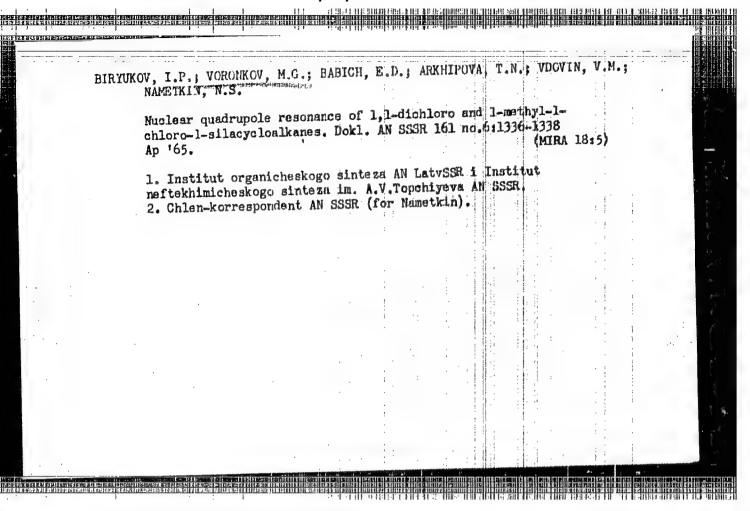


ACCESSION NR: AF5017777  UM/0CB0/65/033/007/1431/1431/1437  AUTHOR: Voronkov, M. G.; Lasskaya, Ye. A.; Pashchinko A. A.  TITLE: Nature of the bonding between water-repeliest draston lices coastings and the surface of materials treated  SOURCE: Zhurnal prikladacy Winni, v. 16, ad. 7, 1765  TOPIC TAGS: hydrophobization, organosilicos competed, water repellent additive, organosilicon coating, polysi oxame film, darbonaulon, glass bond all  ABSTRACT: The fermation of a chemical bond batwaday a water mapel solic polysilotane			
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BIRYUKOV, I.P.; VORONKOV, M.G.; MOTSAREV, G.V.; ROZENDERG, V.R.; SAFIN, I.A.

Muclear quadrupole resonance method of studying organosilicon
compounds containing Si-Cl and C-Cl bonds. Dokl. AN SSSR 162
(MIRA 18:5)
no.1:130-132 Mg. 165.

1. Institut organicheskogo sinteza AN Latviyskoy SSR i Kazanskiy
fiziko-tekhnicheskiy institut AN SSSR. Submitted November 17. 1964.

PART HERBING TO THE STATE OF TH L 17627-66 EWT(m)/EWP(j) ACC NR: AP6001734 SOURCE CODE: UR/0020/65/155/004/0857/0859 Biryukov, I. P.; Voronkov, M. G.; Safin, I. A. AUTHORS: ORG: Institute for Organic Synthesis, Academy of Sciences Latvian SSR (Institut organicheskogo sinteza Akademii nauk LatySSR) TITLE: Correlation of the mean nuclear quadrupole resonants frequency inductive constants of substituents in organylchlorsilands SOURCE: AN SSSR. Doklady, v. 165, no. 4, 1965, 857-859 TOPIC TAGS: NMR, NMR spectroscopy, quadrupole moment, organosilicon compound ABSTRACT: The mean nuclear quadrupole resonance frequency V at 77K. of 36 organylchlorsilanes (general formula RR'R' SiCl35) was correlated with Tait's of the various substituents R, R' and R' on the central silicon atom to extend the work of the authors (Soveshch. po magnitnomu rezonancu, Krain yarak, 1914). The experimental results are presented in graphs and tables. The correlation was effected by the use of equations (1)Card 1/2

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Augule (Lift of the China and China ENT(m)/T/EWP(j) L 32200-66 SOURCE CODE: UR/CO79/66/036/002/031/7/0350 ACC NR: AP6007119 AUTHOR: Orlev, N. F.; Vozenkov, H. G. ORG: Leningrad Institute of Textiles and Light Industry im. S. M. Kirov (Teningradski; institut tekstilinoy i legkoy promyshlennosti); Institute of Organic Synthesis, AN LatSSR(Institut organicheskogo sinteza AN LatSSR) TITIE: Tris(triorganylsilyl) antimonites ( Zhurnal obshchey khimii, v. 36, no. 2, 1966, 347-350 SOURCE: TOPIC TAGS: antimony compound, organic synthetic process, silicon compound ABSTRACT: A study by N. F. Orlov and M. G. Voronkov contains data on two new methods for the synthesis of tris(trialkylsilyl) antimonites. The first method (A), which is simple and economical, consists of reacting a trialkylsilanol with antimony trioxide on heating: 6R<sub>3</sub>SiOH + Sb<sub>2</sub>O<sub>3</sub> → 2Sb(OSiR<sub>3</sub>) + 3H<sub>2</sub>O<sub>4</sub> The water formed is removed by azeotropic distillation with an inert solvent, e.g., benzene, toluene, etc. The yield reaches 70% of the theoretical. Previously, the authors used a similar method to prepare tri-11DC 516.2874516.863

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silanol derivatives of some other trivalent elements or groups: boron, ACC NR: AP6007119 phosphorus, or vanadyl.

The method is similar to that used in the preparation of trialkyl antimonites, Sb(OR)3, by reacting Sb2O3 with alcohols. The higher reactivity of trialkylsilanols, as compared with alcohols, can be explained by their higher acidity caused by the  $p_\pi$  -  $d_\pi$  interactions between silicon and oxygen atoms.

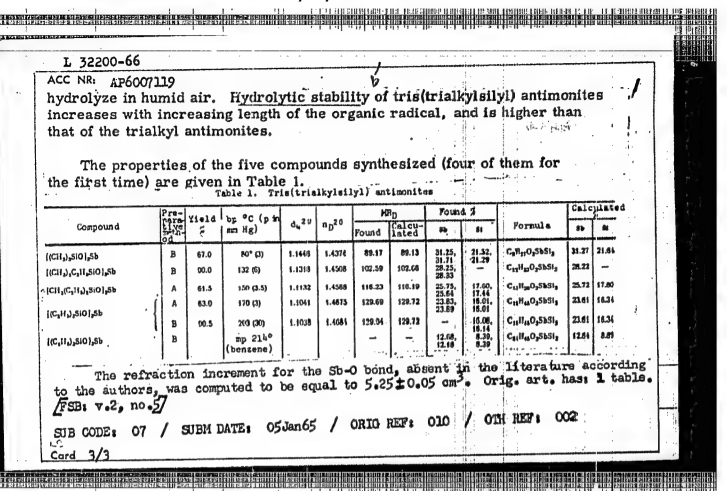
The second method (B) suggested is the silanolysis of trialkyl anti-

monites:

On heating of the mixture the alcohol formed is stripped off. It is advisable to add the silanols in small quantities, e.g., dropwise, especially in benzene solution, in order to prevent silanol condensation, thereby increasing the yield of the desired product.

The presence of sodium in catalytic amounts accelerates the reaction; however, the reaction proceeds readily even without any catalyst,

Tris(trialkylsilyl) antimonites are either colorless liquids or crystals with a weak silanol odor. They are stable on storage in dry air, but Card 2/3



1, 16301-66 EMP(1), EMT(m) SOURCE CODE: UR/C409/66/000/003/0328/0331 AP6023577 ACC NR: AUTHOR: Lukevits, E. Ya.; Voronkov, M. G. ORG: Institute of Organic Synthesis, Academy of Sciences, Latvian SSR, Riga (Institut crganicheskogo sinteza Akademii nauk Latviyskoy SSR) TITLE: Organosilicon compounds of the furan series. Part 12. Furylorganylhydrosilanes SOURCE: Khimiya geterotsiklicheskikh soyedineniy, no. 3, 1966, 328-331 TOPIC TAGS: organosilicon compound, furan compound, silane ABSTRACT: The synthesis of dialkyl(2-furyl)- and dialkyl(5-methyl-2-furyl)hydrosilanes was carried out as follows: B-Li + CHSIR'R' - R- SIHR'R' + LICI.  $R = H_1$   $CH_3$ ;  $R' = CH_3$ ,  $C_2H_5$ ;  $R'' = CH_3$ ,  $C_2H_4$ ,  $n \cdot C_2H_7$ In the reaction of 2-furyllithium with trichlorosilane, the main product is tetra(2furyl)silane (I), and tri(2-furyl)silane is formed in only a slight quantity. 2-furylorganylhydrosilanes react readily with the participation of Si-H bonds, and the reactions are rapid and exothermic. Thus, 2-furyl- and 5-methyl-2-furylorganylhydrosi-547.72+546.2012 Card

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ACC NR: AF6023577

lanes in pyridine instantaneously reduce EgCl2 to Eg metal. In the presence of H2FtCl6, they react vigorously with water dissolved in dioxane. The authors thus synthesized the first representative of 2-furylorganylsilanols, diethyl(2-furyl)silanol:

C2H5

Si-H + H20

C2H5

III

The dohydrocondensation of dimethyl(2-furyl)silane (V) with tetrahydrofurfuryl alcohol is just as rapid, and forms dimethyl(2-furyl)tetrahydrofurfuryloxysilane (VI):

CH3

CH3

CH3

CH3

VI

Addition of III to triethylvinylsilane in the presence of H2FtCl6 also occurs readily:

C2H5

Si-H + CH3=CHSI(C2H3)3 - SI-C3H4SI(C3H3)3

C3H3

VII

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EWT(m)/EWP(j)/T L 40102-66 UR/0080/66/039/006/1345/1351 5/ SOURCE CODE: ACC NR: AP6019567 AUTHOR: Voronkov, M. G.; Pashchenko, A. A.; Lasskaya, Ye. A.; Karibayev, K. K. ORG: Institute of Organic Synthesis, AN LatvSSR (Institut organicheskogo sinteza AN LatvSSR); Kiev Polytechnic Institute (Kiyevskiy politekhnicheskiy institut); Kiev Engineering and Construction Institute (Kiyevekiy inzhenerno-stroitel nyy institut) TITLE: Chemical stability of hydrophobic organosilicon coatings on glass SOURCE: Zhurnal prikladnoy khimii, v. 39, no. 6, 1966, 1345-1351 TOPIC TAGS: polysiloxane, organosilicon compound, protective coating, CHEMICAL STABILITY, CORROSION, CONTED GLASS ABSTRACT: The chemical stability of hydrophobic polyorganosiloxane films deposited on a glass surface from 5% toluene solutions of RSiCl3 was studied by determining their change of wettability, infrared spectra, and thermograms after exposure to the action of aqueous solutions of inorganic acids (HNO3, H2SO4, HC1), bases (NaOH, Ca(OH)2), and salts (Na2SO4, Na2CO3, CaCl2, NaCl, KMnO4). This action was found to break the Si-R bonds. The corrosive attack of the acids and bases increases with their concentration. The greatest stability to the action of corrosive media was displayed by polymethylsiloxane films, and the lowest by polyethylsiloxane ones. Polyallylsiloxanes showed an unexpectedly high chemical stability. Changes in the intensity of the infrared absorption bands and in the heights of exopeaks on the thermograms after exposure to the corrosive agents showed that the stability of the water-661,718,5 UDC: Card 1/2

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repellent films as a function of the organic radical R generally decreases in the

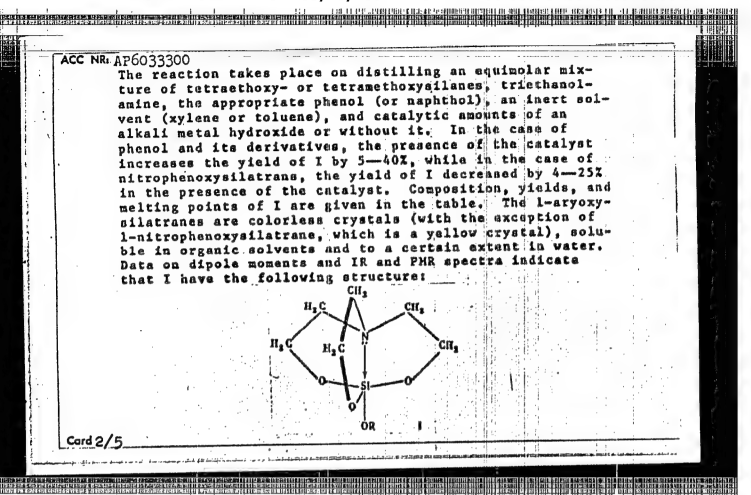
 ${\rm CH_3} > {\rm CH_2}\text{--}{\rm CHCH_2} > {\rm CH_2}\text{--}{\rm CH} > {\rm C_6H_5} > {\rm C_2H_5}$ 

The same order is arrived at by studying the angles of wetting of the polysiloxane films. Orig. art. has: 3 figures and 1 table.

SUB CODE: 07/ SUEM DATE: 26Ju165/ ORIG REF: 012/ OTH REF: 004

Card 2/2 18h

SOURCE CODE: UR/0409/66/000/004/05:1/0514 ACC NR. AP6033300 AUTHOR: Voronkov, M. G.; Zelchan, G. I. ORG: Institute of Organic Synthesis Academy of Sciences LatSSR, Riga (Institut organicheskogo sinteza Akademii nauk LatSSR) Atranes. VIII. 1-Aryloxysilatranes TITLE: SOURCE: Khimiya geterotsiklicheskikh soyedineniy, no. 4, 1966, 511-514 TOPIC TAGS: aryloxysilatrane, biologically compound, heterocyclic base compound, esterification, aromatic hydrocarbon ABSTRACT: The previously unreported 1-aryloxysilatranes (1) ROSI (OCH, CH, ), N (R = aixi). were obtained by the previously described transesterificstion (USSR patent No. 165722, 1964): ROH+Si(OR')4+(HOCH2CH2)3N-ROSi(OCH2CH2)3N+4R'OH, where R=aryl R'=CH3, C2Hs. IDC: 542.95+546.287+547.89 Card 1 /5



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-acc-nrap6033300	2-C, <sub>6</sub> H,	4-02NCH	3-0,NC,H	2-0,NC,H,	2,4,6-C1,C,H,	4-CIC,H	5-CH <sub>3</sub> -	+-{CH,	4.CH,CH	3.CH,C,H,	2-CH <sub>2</sub> C <sub>2</sub> H			 pa s	
		CH.		CH.	1,C,H,	7	5-C113-2-(CH3)2CHC,H3	4-{CH <sub>2</sub> ) <sub>2</sub> CC <sub>2</sub> H <sub>4</sub>	Ŧ	Ŧ			ie.	-Aryloxysila	
	184,5—185,5	182,5—184	197,5—198,5	233—234	230-230,5	166-167	217.5—218,5	252—253	188-189	162,5—163,5	218-219.5	APRIL de	0° °C	trance	
	C14H11NO4SI	C12H11N2O1SI	C12H11N2O,SI	C <sub>12</sub> H <sub>11</sub> N <sub>2</sub> O <sub>3</sub> Si	C;;H;,Cl;NO,Si	C12H14CINO,SI	C,H,NO,SI	C,,H,,NO,Si	ChH,NO,SI	CnH,,NO,SI	C"H"NO'SI	2 L NO 22	Formulas	ROSI(OCH2CH2),N	
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Physiolo	gical activity	of I is con	siderably	lower than	that	
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substitu	ients into arous	atic ring of	the arylo	zysilatrane	18	
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20,, 40	540## 210# 6##			WILLS III.		
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•	5-CH <sub>3</sub> -2-(Cl	p-CIC <sub>8</sub> H <sub>4</sub> 1050 p-GH <sub>3</sub> C <sub>4</sub> H <sub>4</sub> 1275 H <sub>3</sub> ) <sub>2</sub> CHC <sub>6</sub> H <sub>3</sub> 4000				
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SOV/120-59-1-48/50

AUTHORS: Berezin, I. A., Voronkov, M. I.

A Capillary Tap (Kapillyarnyy kran) TITLE:

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PERIODICAL: Pribory i tekhnika eksperimenta, 1959, Nr 1, p 152 (USSR)

ABSTRACT: The device is illustrated diagrammatically in the figure (see lower half of p 152). It is in the form of a two-way valve with two apertures in its plug. The aperture 2 is used in the preliminary evacuation of the tube leading to the gas container. The gas is let in through the capillary tube 1, whose diameter can be arbitrarily small. This permits a very accurate regulation of the pressure of the gas which is admitted into the vacuum system. The paper contains 1 figure.

ASSOCIATION: Fizicheskiky institut AN SSSR (Physics Institute of the Academy of Sciences USSR)

SUBMITTED: January 20, 1958.

Card 1/1

SOV/120-59-1-47/50

AUTHORS: Berezin, I. A., Voronkov, M. I.

A Device for Loading the Samples into a Vacuum System (Bunker dlya zagruzki obraztsov v vakuumnuyu ustanovku)

PERIODICAL: Pribory i tekhnika eksperimenta, 1959, Nr 1, p 152 (USSR)

ABSTRACT: The device described is shown diagrammatically in the figure (see upper half of p 152). The tube 3 contains a polished plug 4 with a helix 2. The tube 3 is inserted into the system in such a way that the open end 1 of the helix is above the crucible of a vacuum furnace. The samples are loaded through the open end of the helical tube in such a way that each turn of the tube contains one sample. The plug is then lubricated with vacuum grease and inserted into the The samples are loaded into the furnace by turning the plug. In this way, the samples are progressively displaced in the helical tube and fall into the oven from the open end of the tube. The paper contains 1 figure.

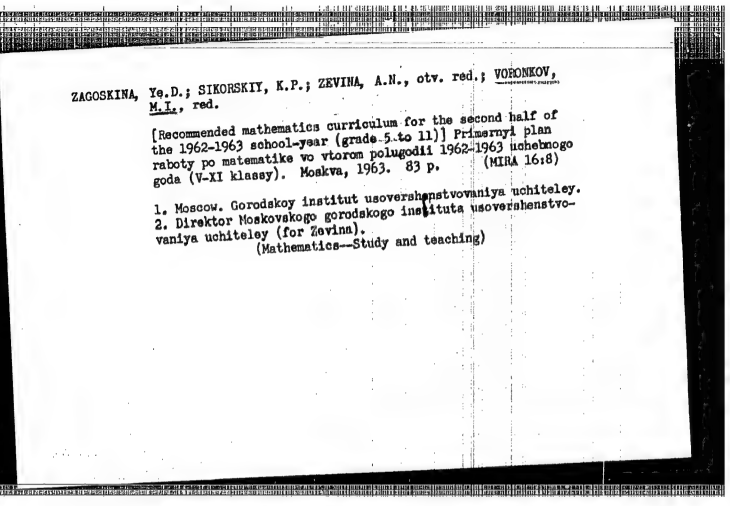
ASSOCIATION: Fizicheskiy institut AN SSSR (Physics Institute of the

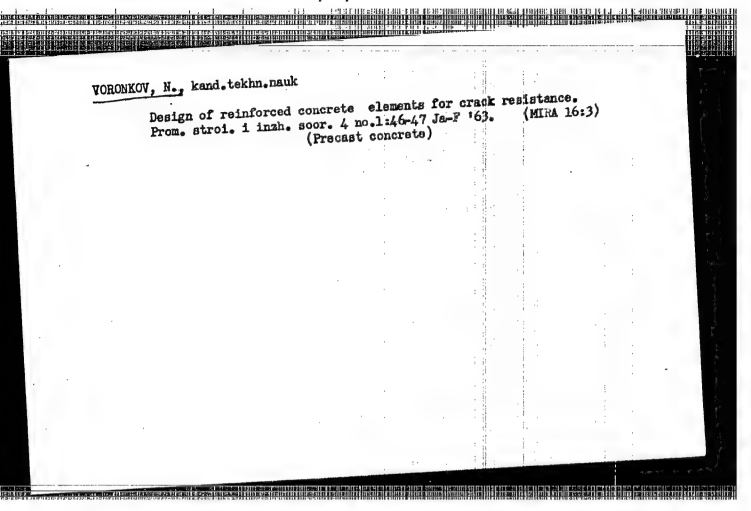
Academy of Sciences, USSR)

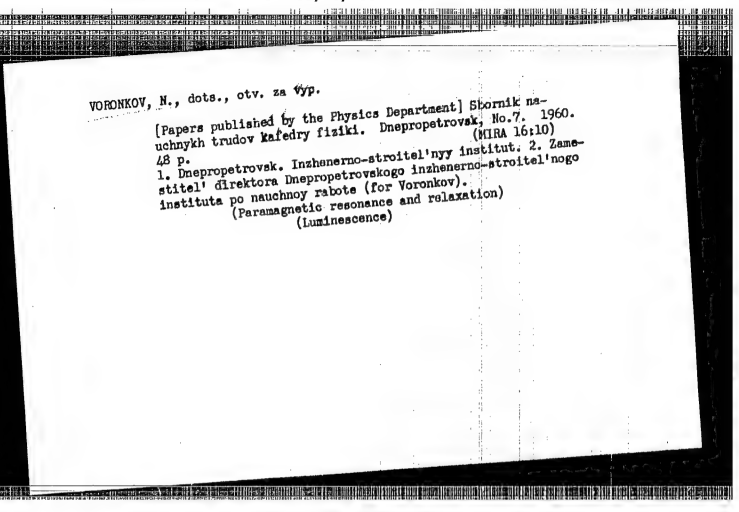
SUBMITTED: January 22, 1958.

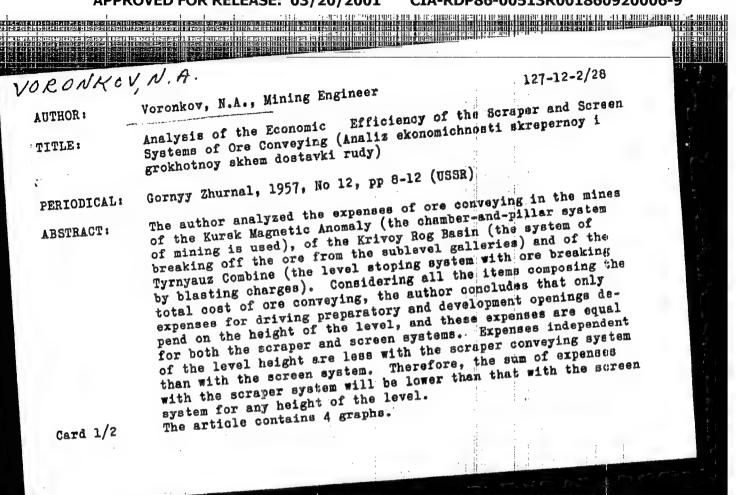
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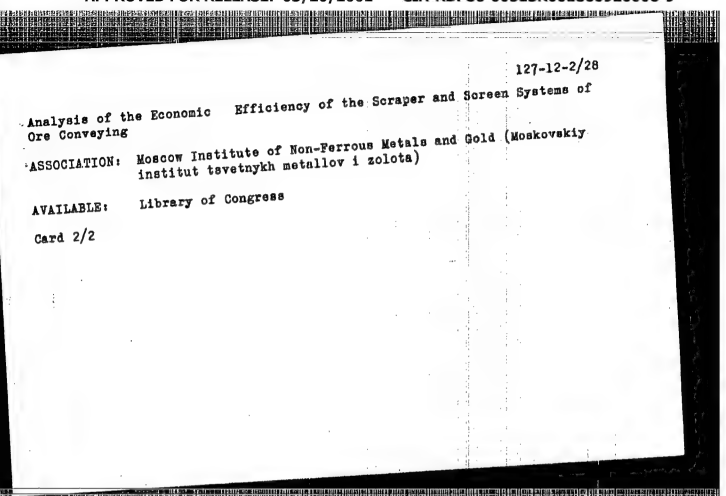
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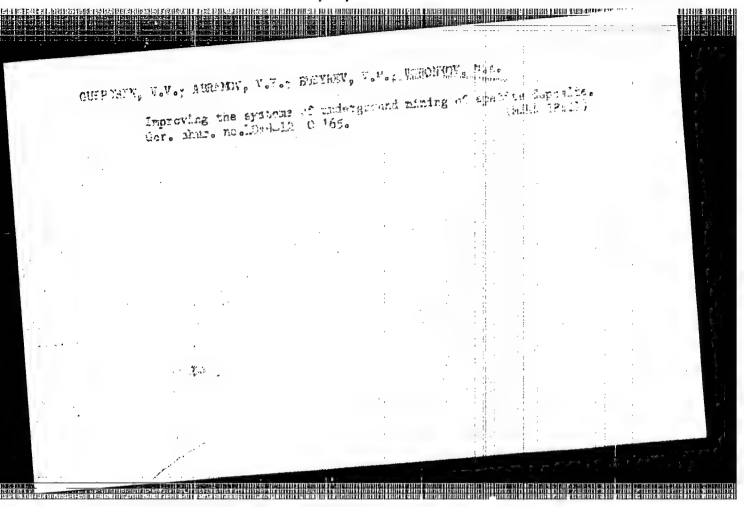


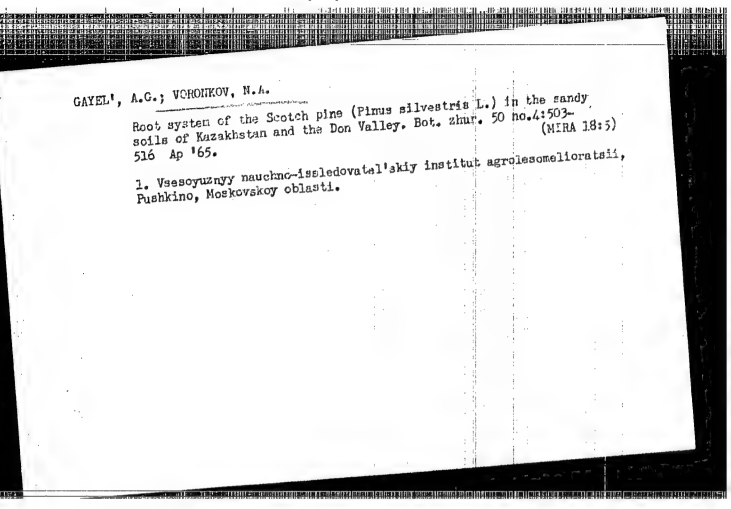


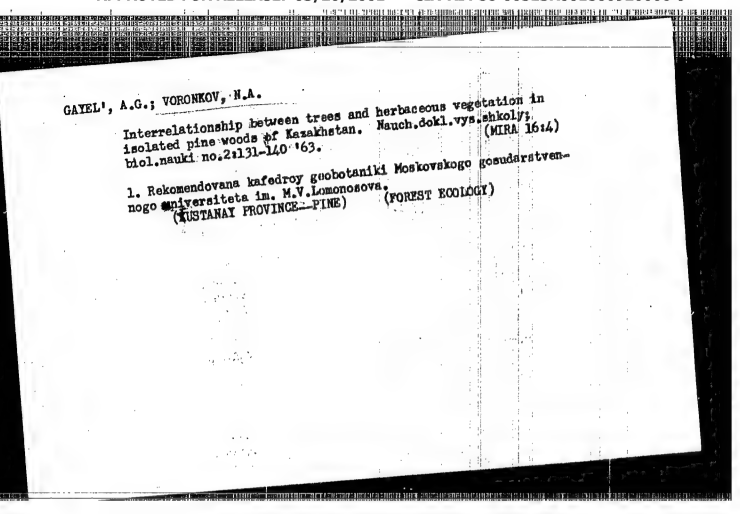


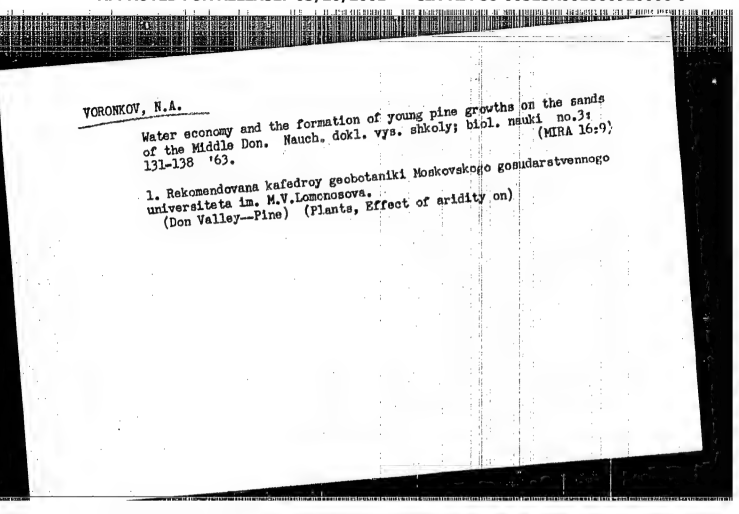


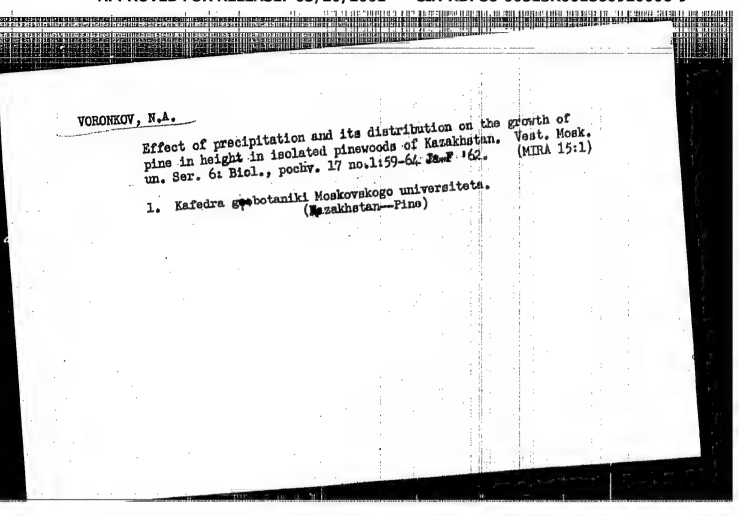


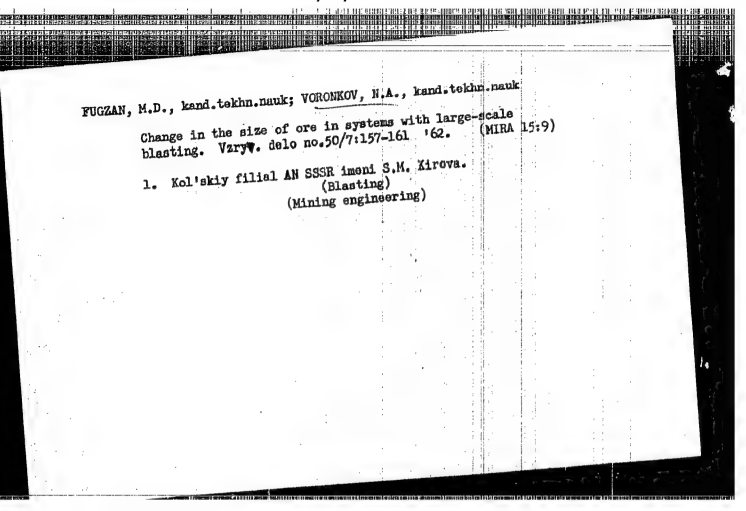


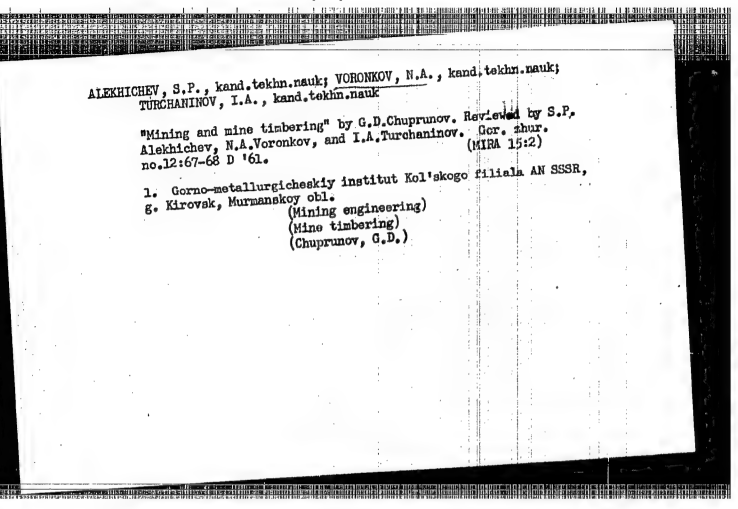


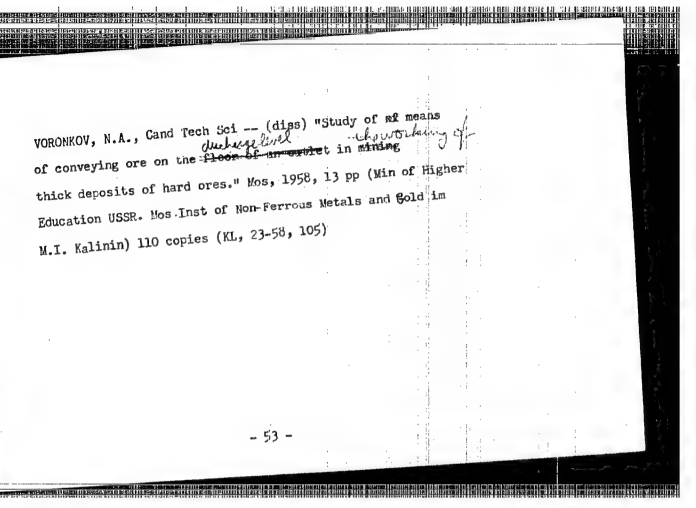












SOV-127-58-10-3/29 -Voronkov, N.A. and Guliy, V.M., Mining Engineers A Comparison of the Scraper and Sifter Methods for Delivery AUTHORS: at the Tekeli Mine (Sravneniye skrepernoy i grokhotnoy TITLE: skhem dostavki na rudnike Tekeli) Gornyy zhurnel, 1958, Nr 10, pp 11-13 (USSR) Research conducted by the authors showed that, under the PERIODICAL: conditions of the Tekeli Mine, where the mining is done by the level caving-in system, the sifter scheme of ore de-ABSTRACT: livery must be preferred to the scraper method. Even though the cost of installation of the sifter method is 80% higher than that of scraper method, it is more advantageous because its productivity is 45% higher. The sifter method also has a lower safety factor than the scraper method. There are 3 tables. card 1/2

